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DIVISIONAL APPLICATION

FOR

NOVEL METHOD FOR PURIFICATION OF RECOMBINANT PROTEINS

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CL-5C1

DESCRIPTION

NOVEL METHOD FOR PURIFICATION OF RECOMBINANT PROTEINS

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Cross-Reference to a Related Application

This application is a continuation-in-part of co-pending application Serial No. 08/698,747, filed August 16, 1996.

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Background of the Invention

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Immobilized metal ion affinity chromatography (IMAC) was first introduced by Porath (Porath, J., J. Carlsson, I. Olsson, G. Belfrage [1975] *Nature* 258:598-599.) under the term metal chelate chromatography and has been previously reviewed in several articles (Porath, J. [1992] *Protein Purification and Expression* 3:263-281; and articles cited therein). The IMAC purification process is based on the employment of a chelating matrix loaded with soft metal ions such as Cu²⁺ and Ni²⁺. Electron-donating groups on the surface of proteins, especially the imidazole side chain of histidine, can bind to the non-coordinated sites of the loaded metal. The interaction between the electron donor group with the metal can be made reversible by lowering the pH or by displacement with imidazole. Thus, a protein possessing electron-donating groups such as histidine can be purified by reversible metal complex/protein interactions.

Several different metal chelating ligands have been employed in IMAC to purify proteins. Iminodiacetic acid (IDA) ligand is a tridentate and thus anchors the metal with only three coordination sites (Porath, J., B. Olin [1983] *Biochemistry* 22:1621-1630). Because of the weak anchoring of the metal, metal leakage has been known to occur. The tris(carboxymethyl)ethylenediamine (TED) ligand is pentadentate and forms a very strong metal-chelator complex. The disadvantage of this is that proteins are bound very weakly since only one valence is left for protein interaction. Nitrilo triacetic acid (NTA) is a tetradentate ligand which attempts to balance the metal anchoring strength with metal-ion

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protein interaction properties (Hochuli, E., H. Dobeli, A. Schacher [1987] J. Chromatography 411:177-184). Other chelating ligands have been reported and are mentioned. See, e.g., Porath (1992), supra. However, these ligands also have certain disadvantages, including decreased bonding capacity, decreased specificity, and increased metal leakage.

In 1991, Ford et al. (Ford, C., I. Suominen, C. Glatz [1991] Protein Expression and Purification 2:95-107) described protein purification using IMAC technology (Ni-NTA ligand) as applied to recombinant proteins having tails with histidine residues (polyhistidine recombinant proteins). This method takes advantage of the fact that two or more histidine residues can cooperate to form very strong metal ion complexes. The NTA chelating ligand immobilized on agarose and loaded with Ni²⁺ has been useful in this method (Hochuli et al., supra; U.S. Patent No. 5,047,513). It is available commercially through Qiagen, Inc. (Chatsworth, CA). However, this resin has the disadvantage that the interchanges between metal ions and poly-histidine recombinant proteins are not optimal. Metal leakage can occur, and background proteins can sometimes contaminate purification of recombinant proteins.

A metal chelating gel, i.e., carboxymethylated aspartate (CM-Asp) agarose complexed with calcium, has been used for purifying native calcium-binding proteins (Mantovaara, T., H. Pertoft, J. Porath [1989] Biotechnology and Applied Biochemistry 11:564-570; Mantovaara, T., H. Pertoft, J. Porath [1991] Biotechnology and Applied Biochemistry 13:315-322; Mantovaara, T., H. Pertoft, J. Porath [1991] Biotechnology and Applied Biochemistry 13:120-126). However, the Ca²⁺-CM-Asp complex described by Mantovaara et al. has among its disadvantages that it does not bind strongly to histidine-tagged recombinant proteins. Another disadvantage, in addition to this inferior binding property, is its non-selectivity for histidine tags.

By contrast, the subject invention comprises the CM-Asp chelating ligand complexed to a transition metal in an octahedral geometry (coordination number of 6). In this unique configuration, the metal complex can be advantageously suited for purification of polyhistidine fused recombinant proteins. This is a novel use of the CM-Asp ligand and is part of the subject of the invention herein described.

Brief Summary of the Invention

The present invention concerns a novel IMAC purification method which employs immobilized carboxymethylated aspartate (CM-Asp) ligands specifically designed for purification of recombinant proteins fused with poly-histidine tags. The new purification method is based upon the CM-Asp chelating matrix having the following structure:

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A general description of the matrix used in the invention and illustrated above is:

When
$$R_4$$
- R_5 - R_6 = H :

M= transition metal ion in a 2+ oxidation state with a coordination number of 6;

 $R_1 = a$ linking arm connecting the nitrogen atom of CM-Asp with R_2 ;

 $R_2 = a$ functional linking group through which CM-Asp linking arm R_1 is connected to R₃;

 R_3 = a polymer matrix, e.g., those polymer matrices typically used in affinity or gel chromatography.

When $R_1 - R_2 - R_3 = H$:

 $R_4 = a$ linking arm connecting the methylene carbon atom of the carboxymethyl group of CM-Asp with R₅;

 $R_5 = a$ functional linking group through which CM-Asp linking arm R_4 is connected to R₆;

 $R_6 = a$ polymer matrix, e.g., those polymer matrices typically used in affinity or gel chromatography.

In a preferred embodiment:

$$M = Fe^{2+}$$
, Co^{2+} , Ni^{2+} , Cu^{2+} , or Zn^{2+} ;

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R_1 = -CH_2CH(OH)CH_2 -, -CH_2(OH)CH_2 - O-CH_2CH(OH)CH_2 -,
      -(CH<sub>2</sub>)<sub>4</sub>NHCH<sub>2</sub>CH(OH)CH<sub>2</sub>-, and -(CH<sub>2</sub>)<sub>2</sub>NHCH<sub>2</sub>CH(OH)CH<sub>2</sub>-;
R_2 = O, S, or NH; and
R_3 = agarose.
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In a particularly preferred embodiment:

 $M = Co^{2+}$;

 $R_1 = CH_2CH(OH)CH_2;$

 $R_2 = O$; and

 R_3 = agarose, cross-linked.

Prior to loading the 6XHis recombinant protein to the resin, recombinant cells can be lysed and sonicated. The lysate can then be equilibrated with an aqueous buffer (pH 8) which itself does not form chelates with the metal. An example of an aqueous that can be used at this step in the described procedure is 50 mM sodium phosphate (pH 8.0)/10 mM Tris-HCl (pH 8.0)/100 mM NaCl, or the like. The equilibration buffer can contain denaturing agents or detergents, e.g., 10% "TRITON X-100," 6 M guanidinium HCl, or the like. After binding the prepared 6XHis recombinant protein on the metal CM-Asp chelating resin (the "CM-Asp resin complex"), the protein-bound resin is washed at pH 7.0 or 8.0. The elution of the protein can be carried out at a constant pH or with a descending pH gradient. In a preferred embodiment, protein elution can be achieved at a pH of about 6.0 to about 6.3. Alternatively, the 6XHis recombinant protein bound to the CM-Asp chelating resin can be washed with low concentrations (less than 100 mM) of imidazole at pH 8.0 and then eluted by increasing the imidazole concentration to 40-100 mM.

Also included as an aspect of the subject invention is a scaled-up synthesis of the CM-Asp derivatized agarose chelating resin. It is an improved version of a previously reported small scale preparation (Mantovaara, T., H. Pertoft, J. Porath [1991] Biotechnology and Applied Biochemistry 13:315-322). The improvement includes particular conditions for oxirane-agarose formation, temperature controlled conjugation of aspartic acid to the oxirane-agarose, and high ionic strength washing to remove extraneously bound metals. These conditions, temperatures, and ionic concentrations are described in detail herein.

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An additional application of the subject invention includes screening for protein function on a microtiter plate or filter. The additional applications for the subject invention also include protein-protein interaction studies, as well as antibody and antigen purification. For example, by immobilization of the Co²⁺ moiety onto 96-well plates by CM-Asp, such plates can be used for quantitation of 6XHistidine-tagged protein, protein-protein interaction studies, diagnostic screening for diseases, antibody screening, antagonist and agonist screening for drugs, and reporter gene assays. Co²⁺ can also be immobilized onto a membrane, e.g., a nylon membrane, by CM-Asp, which can be used to lift proteins from expression libraries to make protein libraries from cells. The membranes also can be used for screening of engineered enzymes. Application of the subject invention can also be extended to purification of any interacting molecule, e.g., nucleic acids or small co-factors.

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Brief Description of the Drawings

Figure 1 is an outline illustrating a process for purifying recombinant 6XHis protein using CM-Asp chelating resin.

Figures 2A-2B show a comparison of Co^{2+} CM-Asp chelating resin with Ni-NTA on 6XHis prepro- α -factor purification under denaturing conditions using pH gradient.

Legend for Figures 2A-2B: lane 1: crude lysate; lane 2: flowthrough; lane 3: washed with 6 M Gu-HCl, 0.1 M NaH₂PO₄, pH 8.0; lane 4: washed with 8 M urea, 0.1 M NaH₂PO₄; lane 5: washed with 8 M urea, 0.1 M NaH₂PO₄, pH 8.0; lane 6: deluted with 8 M urea, 0.1 M NaH₂PO₄, pH 6.3; lane 7: deluted with 8 M urea, 0.1 M NaH₂PO₄, pH 6.3; lane 8: deluted with 8 M urea, 0.1 M NaH₂PO₄, pH 6.3; lane 9: deluted with 8 M urea, 0.1 M NaH₂PO₄, pH 5.9; lane 10: deluted with 8 M urea, 0.1 M NaH₂PO₄, pH 4.5; lane 11: deluted with 6 M Gu-HCl, 0.1 M NaH₂PO₄, 0.2 M acetic acid; lane M: MW size markers.

Figure 3 shows 6XHis tagged DHFR purification by Co²⁺ CM-Asp chelating resin under native conditions. Legend: Lane 1: clarified lysate; lane 2: flowthrough; lane 3: first wash; lane 4: third wash; lane 5: DHFR final elution.

Figure 4 shows 6XHis tagged DHFR purification by Co²⁺ CM-Asp chelating resin under denaturing conditions. Legend: Lane 1: clarified lysate; lane 2: flowthrough; lane

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3: first pH 7.0 wash; lane 4: second pH 7.0 wash; lane 5: DHFR, first pH 6.0 elution; lane6: DHFR, second pH 6.0 elution.

Figure 5 shows 6XHis tagged DHFR purification by Co^{2+} CM-Asp chelating resin under native conditions with increasing concentrations of β -mercaptoethanol. Legend: lane 1: 20 μ l of cell lysate; lanes 2, 4, 6, and 8: 20 μ l of flowthrough; lanes 3, 5, 7, and 9: 5 μ l of eluant.

Figure 6 shows yields of 6XHis DHFR from cell lysates purified by Co^{2+} CM-Asp chelating resin versus Ni-NTA in the presence of β -mercaptoethanol. Protein concentrations were determined by Bradford assay. Yields are expressed as a percentage of total protein in the cell lysate.

Figures 7A-7B show purification of 6XHis GFP by Co²⁺ CM-Asp chelating resin under native conditions. The GFP bands were detected using Clontech's chemiluminescence Western Exposure Kit and overnight exposure to x-ray film.

Legend: lane 1: clarified lysate; lane 2: flowthrough; lane 3: first wash; lane 4: first elution; lane 5: second elution; lane 6: third elution; lane 7: fourth elution.

Figure 8 shows biological activity of 6XHis GFP purified by Co²⁺ CM-Asp chelating resin. Legend: tube 1: cell lysate; tube 3: flowthrough; tube 3: wash; tube 4: first elution; tube 5: second elution; tube 6: third elution.

Detailed Disclosure of the Invention

The subject method, which employs a CM-Asp metal chelating complex, can advantageously be used for purification of recombinant proteins having a polyhistidine tail or "tag."

According to one embodiment of the subject invention, a resin ligand, e.g., CM-Asp, is complexed to a metal other than Ca²⁺, forming a CM-Asp-metal complex. Preferably, the CM-Asp ligand used in the subject invention is complexed with one of the transition metals (known as a third-block transition metal), e.g., Fe²⁺, Co²⁺, Ni²⁺, Cu²⁺, or Zn²⁺ in an octahedral geometry. Other polymer matrices, e.g., polystyrene (as in microtiter plates), nylon (as in nylon filters), SEPHAROSE (Pharmacial, Uppeala, Sweden) or the like, can be used with

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the subject invention and would readily be recognized by persons of ordinary skill in the art. The poly-histidine tag possesses "neighboring" histidine residues which can advantageously allow the recombinant protein to bind to these transition metals in a cooperative manner to form very strong metal ion complexes. This cooperative binding refers to what is commonly known in the art as a "neighboring histidine effect." For purposes of the subject invention, and as would be understood by a person of ordinary skill in the art, a "strong" or "very strong" metal ion complex refers to the bond strength between the metal ion and the chelating ligand. A strong or very strong metal ion complex, for example, allows little or essentially no metal leakage from the complex so that the purified protein, e.g., a recombinant protein having a polyhistidine tag, is not contaminated with undesired or "background" protein from a mixture being purified.

The CM-Asp metal complex offers two available valencies that can form strong and reversible metal complexes with two adjacent histidine residues on the surface of the recombinant protein. Another advantage to using the CM-Asp ligand is its ability to strongly anchor the metal ion whereby metal ion leaking can be virtually eliminated compared to metal leakage observed for other complex binding agents, e.g., Ni-NTA. In a more preferred embodiment, Co^{2+} can be used as the transition metal with CM-Asp. The Co^{2+} -CM-Asp can be less sensitive to reducing agents, such as β -mercaptoethanol. Metal ion leakage has been shown to remain low, even negligible, in the presence of up to 30 mM β -mercaptoethanol.

One embodiment of the purification process of the subject invention is as follows:

- 1. Prepare lysate/sonicate containing recombinant 6XHis protein according to standard procedures and techniques well known in the art.
- 2. Bind 6XHis protein onto metal-loaded CM-Asp chelating resin at slightly basic pH, e.g., about pH 8.0.
- 3. Wash protein/resin complex at the same basic pH (about pH 8.0). Optional washes at a pH of about 7.0 or with imidazole additive can also be included.
- 4. Elute pure recombinant 6XHis protein with an elution buffer having a pH of about 6.0-6.3 or, in the alternative, an elution buffer having a pH of about 8.0, plus about 40 to about 100 mM imidazole.

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The steps involved in a preferred embodiment of the purification process of the subject invention are illustrated in Figure 1. The subject process can be employed batchwise, in spin columns, and in large-scale continuous-flow columns.

Buffers used in the above procedures are standard buffers typically used in similar procedures, with appropriate adjustments and modifications made as understood in the art. For example, a high ionic strength buffer, e.g., 50 mM phosphate/10 mM Tris/100 mM NaCl can be used, with the pH adjusted as needed. The phosphate salt component can range from a concentration of 10-100 mM; Tris from 5-25 mM; and NaCl from 50-200 mM.

Optimal elution conditions depend on the type of impurities, the amount of protein to be purified, and unique properties of the protein, and are determined on a case-by-case basis as would be readily recognized by a person of ordinary skill in the art.

The subject invention also pertains to a method for synthesizing carboxymethylated aspartate chelating matrices, comprising the steps of:

- (a) Michael addition of the α -amino function of monoprotected α,ω -diamino acids to maleic acid;
- (b) deprotecting the ω -amino functionality; and
- (c) attaching the chelator primary amine molecule to a solid matrix.

Following are examples which illustrate procedures for practicing the invention. These examples should not be construed as limiting. All percentages are by weight and all solvent mixture proportions are by volume unless otherwise noted.

Example 1 - Large-Scale Preparation of CM-Asp Chelating Resin

SEPHAROSE CL-6B or CL-6B (Pharmacia, 8.0 L) is washed with ddH₂O, suction dried, and transferred to a 22-L round bottom flask equipped with mechanical stirring. Epichlorohydrin (about 2.0 L) is added, the Sepharose resin mixed to a thick suspension, and allowed to stand at room temperature for about 20 minutes. A solution of sodium hydroxide (about 560 g) and sodium borohydride (about 48 g) in approximately 6400 mL ddH₂O is added and the mixture stirred overnight at ambient temperature. The oxirane-derivatized

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resin, collected by filtration, is washed ten times with ddH₂O (about 10 L each), once with 10% sodium carbonate (about 10 L), suction dried, and transferred to a 22L round bottom flask. A specimen of the oxirane derivatized resin treated with 1.3 M sodium thiosulfate is titrated to approximately pH 7.0 to determine the oxirane concentration (preferably, >700 µmol/g).

To a solution of sodium hydroxide (approximately 268 g) in about 7.6 L ddH₂O is added L-aspartic acid (about 575 g) and sodium carbonate (about 1700 g), keeping the temperature below about 25 °C. The pH is adjusted to approximately 11.0 and the solution added to the resin. Using mechanical stirring and a heating mantle, the reaction mixture is brought to about 80 °C for 4 hours and allowed to cool to room temperature overnight. The resin was collected by filtration, washed ten times with ddH₂O (about 10 L each), once with 10% sodium carbonate (about 10 L), suction dried, and transferred to a 22-L round bottom flask equipped with mechanical stirring.

To an ice-cooled solution of sodium hydroxide (about 900 g) in 12 L ddH₂O was added bromoacetic acid (about 3000 g) in approximately 750 g increments, keeping temperature below about 30°C. Sodium carbonate (about 660 g) is added and the pH is adjusted to about 10. The resin is reacted with the solution at ambient temperature overnight. The resin is collected by filtration, washed six times with ddH₂O (about 10 L each), six times with 10% acetic acid, and ten times with ddH₂O. Washing is continued with ddH₂O until the pH reached about 6.0 by litmus paper. The CM-Asp chelating resin was suction dried in preparation for metal loading.

Example 2 - Preparation of C-Linked CM-Asp Chelating Resin

No-Carbobenzyloxy-L-lysine (6.15 g) and excess maleic acid (17.6 g) were dissolved in 2 M NaOH (35 mL). The solution was refluxed for 24 hr and allowed to cool to ambient. The pH was adjusted to 3 with 6 M HCl and chilled. The mixture was filtered to remove the unreacted maleic acid which was washed with water (15 mL). The filtrate and washings were combined and evaporated. The residue was dissolved in 4% ammonium formate (120 mL) and degassed briefly. 10% Pd/C (600 mg) was added and the mixture was refluxed

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under Ar with stirring for 5 hr. The mixture was filtered through celite and the filtrate evaporated. The residue was dissolved in a solution of sodium hydroxide (1.3 g) and sodium carbonate (8.7 g) in dd H₂O (40 mL). The final pH was adjusted to 11. This solution is added to the oxirane-derivatized resin (40 mL bed volume) prepared as described in Example 1. The mixture was stirred at 60° with mechanical stirring for 4 hr and at ambient for 16 hr. The resin was collected by filtration, washed with ddH₂O (6 x 100 mL), 10% HOAc (6 x 100 mL), and ddH₂O (6 X 100 mL) or until the pH reached about 6 by litmus paper test. The C-Linked CM-Asp chelating resin was suction dried in preparation for metal loading.

Example 3 - Metal Loading of CM-Asp Chelating Resin

The CM-Asp chelating resin of Example 1 (about 1 L of suction dried bed volume) is treated with a transition metal ion solution, e.g., 2 L of either 200 mM of cobalt chloride hexahydrate, nickel sulfate hexahydrate, copper sulfate pentahydrate, or zinc chloride, according to the metal ion deserved. The resin is reacted with the 200 mM metal solution at room temperature for approximately 72 hours and then collected by filtration. The metal loaded CM-Asp chelating resin is washed five times with ddH₂O (about 1 L each), two times with 100 mM NaCl (about 1 L each), six times with ddH₂O (about 1 L each), and once with 20% aq. ethanol (about 1 L). The resin can be stored in 20% aq. ethanol.

Example 4 – Comparison of Co²⁺ CM-Asp Resin With Ni²⁺-NTA on Recombinant 6XHis Prepro-α-Factor Under Denaturing Conditions Using pH Gradient

For a qualitative comparison of the purification of Co²⁺ CM-Asp chelating resin and Ni-NTA under denaturing conditions, the C-terminal, 6XHis-tagged prepro-α-factor of S. cerevisiae was expressed in E. coli. One gram bacterial cell pellet was lysed in 6 M guanidinium-HCl (Gu-HCl) and 0.1 M NaH₂PO₄, pH 8.0. Three milliliters of clarified lysate was loaded onto a Co²⁺ CM-Asp chelating resin gravity flow column. The resin-proteins mixture was washed with 8 M urea, 0.1 M NaH₂PO₄, pH 8.0, and deluted with 8 M urea, 0.1 M NaH₂PO₄ at three different pHs, 6.3, 5.9, and 4.5. Finally, all bound proteins were deluted with 6 M Gu-HCl, 0.1 M NaH₂PO₄, 0.2 M acetic acid. Samples from each step were loaded

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on a 12% polyacrylamide/SDS gel, electrophoresed, and the gel was stained with Coomassie blue. The 6XHis-tagged prepro- α -factor was deluted at pH 6.3 as a single prominent band on the gel.

In the same manner, 3 ml of clarified lysate was loaded onto a Ni-NTA gravity flow column. The resin-proteins mixture was washed and deluted the same as above. Samples from each step were loaded on a 12% polyacrylamide/SDS gel, electrophoresed, and the gel was stained with Coomassie blue. There were more than 10 protein bands in elution at pH 6.3. The 6XHis-tagged prepro- α -factor was a minor band among them. The majority of the protein was deluted at pH 4.5 without any other contaminant proteins. This demonstrated that the highly purified 6XHis-tagged prepro- α -factor was deluted from Co²⁺ CM-Asp chelating resin at the conditions (pH 6.3) under which Ni-NTA was still releasing contaminants. The affinity of Co²⁺ CM-Asp chelating resin to 6XHis-tagged prepro- α -factor was more selective than Ni-NTA to the protein.

Results show that highly purified 6XHis-tagged protein elutes from Co²⁺ CM-Asp chelating resin while Ni-NTA is still releasing contaminants. See Figures 2A-2B: Figure 2A: Results after using 1 ml of Co²⁺ CM-Asp chelating resin. Figure 2B: Results after using 1 ml of nickel-NTA.

Example 5 – Recombinant 6XHis DHFR Purification with CM-Asp Resin Under Native Conditions

N-terminal, 6XHis-tagged mouse dihydrofolate reductase (DHFR, MW 20.3 kDa) was expressed in *E. coli* cells. Cells were then pretreated with 0.75 mg/ml lysozyme and disrupted in lysis buffer (100 mM NaH₂PO₄, 10 mM Tris-HCl, pH 8.0) by mechanical shearing, 800 µl of the clarified lysate was applied to 100 µl of Co²⁺ CM-Asp chelating resin, pre-equilibrated with lysis buffer, and washed with one ml of lysis buffer three times. All bound protein was deluted by 300 µl of 100 mM EDTA, pH 8.0. Twenty microliters of lysate and 40 µl of each subsequent fraction from elution were run on a 12% polyacrylamide/SDS gel. The gel was stained with Coomassie blue. One single protein band was shown at a position of MW 20.3 kDa. Results showed the selective binding

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affinity of Co2+ CM-Asp chelating resin to 6XHis-tagged DHFR under native purification conditions. No discernable binding of host proteins occurred.

Results show that Co²⁺ CM-Asp chelating resin has selective binding affinity for 6X Histidines. No discernable binding of host proteins occurred. See Figure 3.

Example 6 - Recombinant 6XHis DHFR Purification with CM-Asp Resin Under Denaturing Conditions

N-terminal 6XHis-tagged mouse DHFR was expressed in a 25-ml culture of E. coli. Cells were pelleted, resuspended in lysis buffer (100 mM NaH₂PO₄, 10 mM Tris-HCl, 8 M urea, pH 8.0), and disrupted by vortexing. Six hundred microliters of clarified lysate were applied to a Co²⁺ CM-Asp chelating resin spin column containing 0.5 ml of Co²⁺ CM-Asp chelating resin-NX metal affinity resin and centrifuged for 2 minutes at 2,000 x g. The column was washed twice with 1 ml of wash buffer (100 mM NaH₂PO₄, 10 mM PIPES, pH 7.0), and bound proteins were deluted with 600 µl of elution buffer (20 mM PIPES, 100 mM NaCl, 8 M urea, pH 6.0). Twenty microliters of lysate and 40 µl of each subsequent fraction from the elution were loaded onto a 12% polyacrylamide/SDS gel and electrophoresed. The gel was stained with Coomassie blue. One single protein band was shown at the position of 20.3 kDa. Results showed the selective binding affinity of Co²⁺ CM-Asp chelating resin to 6XHis-tagged DHFR under denaturing conditions. The binding properties of Co²⁺ CM-Asp chelating resin to 6X histidines allow proteins deluted under mild pH conditions (pH 6.0) that protect protein integrity.

Results show that bound protein can be deluted at mild pH (pH 6.0). This indicates that the binding properties of Co²⁺ CM-Asp chelating resin allow protein elution under mild pH conditions that protect protein integrity. See Figure 4.

Example 7 - Recombinant 6XHis DHFR Purification with CM-Asp Resin Under Native Conditions with Increasing Concentrations of Beta-Mercaptoethanol

N-terminal, 6XHis-tagged mouse DHFR was expressed in E. coli. Twenty-five milliliters of cell culture were disrupted in 2 ml of sonication buffer (100 mM NaH₂PO₄, 10

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mM Tris-HCl, and 100 mM NaCl, pH 8.0) by freezing and thawing. Then, 2.66 ml of clarified lysate were applied to a 200- μ l Co²⁺ CM-Asp chelating resin gravity flow column, pre-equilibrated with the sonication buffer. The proteins/resin mixtures were washed three times with sonication buffer, pH 8.0. All bound proteins were deluted with 600 μ l of 100 mM EDTA, pH 8.0. To test the effect of β -mercaptoethanol on the Co²⁺ CM-Asp chelating resin purification under native conditions, all buffers used here contained either 0, 10, 20, or 30 mM β -mercaptoethanol. Samples from each elution were electrophoresed on a 12% polyacrylamide/SDS gel, and the gel was stained with Coomassie blue. One single protein band at the position of MW 20.3 kDa was shown from all elutions. The presence of β -mercaptoethanol did not obsolete the purity of 6XHis-tagged DHFR purified by Co²⁺ CM-Asp chelating resin. With up to 30 mM β -mercaptoethanol in all purification buffers, there was no predominant band at 20.3 kDa in flowthroughs, indicating that no loss of metal occurred during protein purification using Co²⁺ CM-Asp chelating resin in the presence of β -mercaptoethanol.

Results show that with up to 30 mM β -mercaptoethanol in the purification buffer, there is no predominant band at 20.3 kDa in the flowthrough, indicating that no loss of metal occurred during protein purification using Co²⁺ CM-Asp chelating resin in the presence of β -mercaptoethanol. See Figure 5.

Example 8 — Yields of 6XHis DHFR From Cell Lysates Purified by CM-Asp Versus Ni-NTA in the Presence of Beta-Mercaptoethanol

N-terminal, 6XHis-tagged DHFR was expressed and purified by Co^{2+} CM-Asp chelating resin under native conditions as described in Example 7. Protein concentrations were determined by Bradford assay. Yields were expressed as a percentage of total protein in the cell lysate. The yields of purified 6XHis-tagged DHFR were 14%, 28%, 34%, and 35% respectively, with β -mercaptoethanol present in purification buffers at the concentrations of 0, 10, 20, and 30 mM. The protein was purified by Ni-NTA under the same native conditions; the yields of purified 6XHis-tagged DHFR were 4%, 8.8%, 3.4%, and 4% respectively with β -mercaptoethanol present in purification buffers at the

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concentrations of 0, 10, 20, and 30 mM. The yields of purified 6XHis-tagged DHFR were significantly higher when using Co^{2+} CM-Asp chelating resin compared to Ni-NTA under native purification conditions with β -mercaptoethanol. This indicates that the metal ion on Co^{2+} CM-Asp chelating resin is strongly anchored to SEPHAROSE beads by a CM-ASP metal chelator that is ideal for binding octahedral metals.

Results show that the yields of purified 6XHis DHFR are significantly higher at 10, 20, and 30 mM β -mercaptoethanol using Co²⁺ CM-Asp chelating resin compared to using Ni-NTA. This indicates that the metal ion on Co²⁺ CM-Asp chelating resin is strongly anchored to sepharose beads by CM-Asp metal chelator that is advantageous for binding octahedral metals. See Figure 6.

Example 9 – Purification of 6XHis GFP by Co²⁺-CM-Asp Under Native Conditions

N-terminal, 6XHis-tagged green fluorescent protein (GFP) was expressed in *E. coli* cells. Cells were pelleted, resuspended in sonication buffer (100 mM NaH₂PO₄, 10 mM Tris-HCl, and 100 mM NaCl, pH 8.0), and disrupted by freezing and thawing three times. Two milliliters of clarified lysate were applied to 400 µl of Co²⁺ CM-Asp chelating resin, preequilibrated with sonication buffer, and washed three times with 2 ml of sonication buffer, pH 8.0. The 6XHis-tagged GFP was deluted with 400 µl of 75 mM imidazole buffer containing 20 mM Tris-HCl and 100 mM NaCl, pH 8.0. Samples from each purification step were loaded onto a 12% polyacrylamide/SDS gel, electrophoresed, and the gel was stained with Coomassie blue. One single band was shown at the position of MW 27.8 kDa in the elution with 75 mM imidazole. This demonstrated that 6XHis-tagged GFP selectively bound on Co²⁺ CM-Asp chelating resin and can be deluted with low concentration of imidazole under native purification conditions.

Samples from each purification step were also loaded on a 12% polyacrylamide/SDS gel, electrophoresed, and transblotted to a PVDF membrane. The proteins on the blot were probed with anti-GFP monoclonal antibody. One single GFP band was clearly shown in the samples of cell lysate and elution. There was no GFP band shown in flowthrough, which

indicated that all expressed GFP in cell lysate was bound to $\mathrm{Co^{2+}}$ CM-Asp chelating resin.

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Results show that (Figure 7A) Coomassie blue stained gel shows one single band in 75 mM imidazole elution. This indicates that 6X Histidines selectively bound on Co²⁺ CM-Asp chelating resin. Western analysis data shows no GFP in flowthrough which indicates the high affinity between Co²⁺ CM-Asp chelating resin and 6X histidines (Figure 7B).

Example 10 - Biological Activity of 6XHis GFP Purified by Co²⁺ CM-Asp

N-terminal, 6XHis-tagged GFP was expressed in *E. coli*. Cell lysate was prepared as described in Example 7. The cell lysate was loaded onto a 2-ml Co²⁺ CM-Asp chelating resin Disposable Gravity Column, and purified using the Batch/Gravity Flow column purification method as described in Example 9. The column was washed with sonication buffer three times and deluted with 100 mM EDTA, pH 8.0. Samples were collected in microfuge tubes from each purification step. The fluorescence of all collected samples was detected using an UltraLum Electronic U.V. Transilluminator. Samples of cell lysate and elution showed strong fluorescence. This experiment demonstrated that 6XHis-tagged GFP can be purified to homogeneity by Co²⁺ CM-Asp chelating resin under native conditions and maintains biological activity.

The photo of samples from each purification step shows that GFP can be purified to homogeneity by Co²⁺ CM-Asp chelating resin under native conditions, and the fluorescence indicates that GFP purified by Co²⁺ CM-Asp chelating resin still maintains its biological activity. See Figure 8.

It should be understood that the examples and embodiments described herein are for illustrative purposes only and that various modifications or changes in light thereof will be suggested to persons skilled in the art and are to be included within the spirit and purview of this application and the scope of the appended claims.